Refine Search

Search Results -

Terms	Documents
L13 and (SPD or MBL\$ or SPA or collectin-43)	31

US Pre-Grant Publication Full-Text Database

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Search:

L15			Refine Search
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Search History

DATE: Thursday, October 19, 2006 Purge Queries Printable Copy Create Case

Set Name side by side	Query	Hit Count	<u>Set</u> <u>Name</u> result set
DB=U	USPT; PLUR=YES; OP=OR		
<u>L15</u>	L13 and (SPD or MBL\$ or SPA or collectin-43)	31	<u>L15</u>
<u>L14</u>	L13 and collectin\$	29	<u>L14</u>
<u>L13</u>	17 and chimeric	203	<u>L13</u>
<u>L12</u>	17 with chimeric	0	<u>L12</u>
<u>L11</u>	17 and collectins	2	<u>L11</u>
<u>L10</u>	L8 and (fusion or heterologous)	81	<u>L10</u>
<u>L9</u> -	L8 with (fusion or heterologous)	0	<u>L9</u>
<u>L8</u>	L7 with ligand	82	<u>L8</u>
<u>L7</u>	(tumor adj necrosis adj factor adj superfamily) or ((tumor adj necrosis adj factor) with superfamily)	259	<u>L7</u>
<u>L6</u>	L5 and ligand	· 9	<u>L6</u>
<u>L5</u>	(tumor adj necrosis adj factor adj superfamily)	10	<u>L5</u>
<u>L4</u>	TNFSF	0	<u>L4</u>

<u>L3</u>	TNFSF adj ligand	0	<u>L3</u>
<u>L2</u>	L1 with TNFSF	0	<u>L2</u>
<u>L1</u>	extracellualr adj domain	4	<u>L1</u>

END OF SEARCH HISTORY

FILE 'HOME' ENTERED AT 11:26:39 ON 19 OCT 2006

=> file medline caplus biosis embase uspatful COST IN U.S. DOLLARS SINCE FILE TOTAL

ENTRY

SESSION FULL ESTIMATED COST 0.21 0.21

FILE 'MEDLINE' ENTERED AT 11:27:24 ON 19 OCT 2006

FILE 'CAPLUS' ENTERED AT 11:27:24 ON 19 OCT 2006
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FILE 'USPATFULL' ENTERED AT 11:27:24 ON 19 OCT 2006 CA INDEXING COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

=> s collectin? (W) (fusion or chimeric)
L1 13 COLLECTIN? (W) (FUSION OR CHIMERIC)

=> s |1 and TNFSF L2 2 L1 AND TNFSF

=> duplicate remove I2

DUPLICATE PREFERENCE IS 'CAPLUS, USPATFULL'
KEEP DUPLICATES FROM MORE THAN ONE FILE? Y/(N):n
PROCESSING COMPLETED FOR L2
L3 2 DUPLICATE REMOVE L2 (0 DUPLICATES REMOVED)

=> d I3 1- ibib, abs YOU HAVE REQUESTED DATA FROM 2 ANSWERS - CONTINUE? Y/(N):y

L3 ANSWER 1 OF 2 USPATFULL on STN ACCESSION NUMBER: 2005:183444 USPATFULL <<LOGINID::20061019>> TITLE: Multimeric fusion proteins of TNF superfamily ligands INVENTOR(S): Kornbluth, Richard S., La Jolla, CA, UNITED STATES

NUMBER KIND

DATE

PATENT INFORMATION: US
2005158831 A1 20050721
APPLICATION INFO.: US 2005-87348
A1 20050322 (11)
RELATED APPLN. INFO.: Continuation of
Ser. No. US 1999-454223, filed on 9 Dec
1999, PENDING

NUMBER DATE

PRIORITY INFORMATION: US 1998111471P 19981209 (60)
DOCUMENT TYPE: Utility
FILE SEGMENT: APPLICATION
LEGAL REPRESENTATIVE: Lisa A. Haile,
J.D., Ph.D., DLA PIPER RUDNICK GRAY
CARY

US LLP, Suite 1100, 4365

Executive Drive, San Diego,

CA, 92121-2133, US

NUMBER OF CLAIMS: 24
EXEMPLARY CLAIM: 1-15

NUMBER OF DRAWINGS: 7 Drawing

Page(s)

LINE COUNT: 1657

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A method for constructing stable bioactive fusion proteins of the

difficult to express turn or necrosis factor superfamily (***TNFSF***

), and particularly members CD40L (CD 154) and RANKL/TRANCE, with

collectins, particularly pulmonary surfactant protein D (SPD) is described. Single trimers of these proteins lack the full stimulatory efficacy of the natural membrane forms of these proteins in many cases. The multimeric nature of these soluble fusion proteins enables them to engage multiple receptors on the responding cells, thereby, mimicking the effects of the membrane forms of these ligands. For CD40L-SPD, the resulting protein stimulates B cells, macrophages, and dendritic cells, indicating its potential usefulness as a vaccine adjuvant. The large size of these fusion proteins makes them less likely to diffuse into the circulation, thereby limiting their potential systemic toxicity. This property may be especially useful when these proteins are injected locally as a vaccine adjuvant or tumor immunotherapy agent to prevent them from diffusing away. In addition, these and other ***TNFSF*** -***collectin*** ***fusion*** proteins present new possibilities for the expression of highly active, multimeric, soluble ***TNFSF*** members.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L3 ANSWER 2 OF 2 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2001:435124 CAPLUS <<LOGINID::20061019>> DOCUMENT NUMBER: 135:45182 TITLE: Multimeric forms of TNF superfamily ligands INVENTOR(S): Kornbluth, Richard S. PATENT ASSIGNEE(S): USA SOURCE: PCT Int. Appl., 73 pp. CODEN: PIXXD2 **DOCUMENT TYPE:** Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

KIND DATE

DATE

PATENT NO.

APPLICATION NO.

W: AU, CA, JP RW: AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE CA 2393659 AA 20010614 CA 2000-2393659 20000320 A1 20020904 EP EP 1235853 20000320 2000-919485 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, FI, CY US 2005158831 A1 20050721 US 2005-87348 20050322 PRIORITY APPLN. INFO.: US 1999-454223 A 19991209 US 1998-111471P P 19981209 WO 2000-W 20000320 US7380 AB A method for constructing stable bioactive fusion proteins of the difficult to express tumor necrosis factor superfamily (***TNFSF***), and particularly members CD40L (CD154) and RANKL/TRANCE, with collectins, particularly pulmonary surfactant protein D (SPD) is described. Single trimers of these proteins lack the full stimulatory efficacy of the natural membrane forms of these proteins in many cases. The multimeric nature of these sol. fusion proteins enables them to engage multiple receptors on the responding cells, thereby, mimicking the effects of the membrane forms of these ligands. For CD40L-SPD, the resulting protein stimulates B cells, macrophages, and dendritic cells, indicating its potential usefulness as a vaccine adjuvant. The large size of these fusion proteins makes them less likely to diffuse into the circulation, thereby limiting their potential systemic toxicity. This property may be esp. useful when these proteins are injected locally as a vaccine adjuvant or tumor immunotherapy agent to prevent them from diffusing away. In addn., these and other ***TNFSF*** -***collecting*** ***fusion*** proteins present new possibilities for the expression of highly active,

WO 2001042298

WO 2000-US7380

A1 20010614

20000320

multimeric, sol. ***TNFSF*** members. REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> d l1 1- ibib, abs YOU HAVE REQUESTED DATA FROM 13 ANSWERS - CONTINUE? Y/(N):y

L1 ANSWER 1 OF 13 CAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2004:252645 CAPLUS <<LOGINID::20061019>> DOCUMENT NUMBER: 140:286164 TITLE: Fusion proteins of complement activating proteins and lectins for lectin-mediated activation of complement INVENTOR(S): Kongerslev, Leif;

Weilguny, Dietmar; Matthiesen, Finn PATENT ASSIGNEE(S): Natlmmune A/S, Den. SOURCE: PCT Int. Appl., 127 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

WO 2004024925 A2 20040325 WO 2003-DK585 20030910 WO 2004024925 C1 20040521 A3 20040624 WO 2004024925 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,

KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG AU 2003260286 A1 20040430 AU 2003-260286 20030910 A2 20050615 EP EP 1539964 2003-794818 20030910 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK CN 1694961 Α 20051109 CN 20030910 2003-825026 JP 2005537807 T2 20051215 JP 2004-535018 20030910 US 2006188963 A1 20060824 US 2005-527191 20050310 PRIORITY APPLN. INFO.: DK A 20020910 2002-1328 WO 2003-

DK585 W 20030910 AB Fusion proteins of complement activating proteins that can be used to stimulate the lectin-dependent pathway of complement activation in improving the response to infection are described. The proteins are fusion products of complement-activating proteins and lectins such as collectins, L-ficolin, or mannan-binding lectins. These fusion proteins are suitable for therapeutic reconstitution or improvement of opsonic or bactericidal activity of the complement system, i.e. enhancing the ability of the immune defense to recognize and kill microbial pathogens, and accordingly, the invention relates to a medicament comprising the fusion protein, methods for producing said fusion protein and methods for treating diseases, in particular infections.

L1 ANSWER 2 OF 13 CAPLUS COPYRIGHT 2006 ACS on STN **ACCESSION NUMBER:** 2004:183045 CAPLUS <<LOGINID::20061019>> **DOCUMENT NUMBER:** 140:234386 TITLE: Chimeric proteins comprising lectin carbohydrate-binding

domain and cell surface protein

ligand for modulating	US 2004122217 A1 20040624
immune response to antigen	US 2003-666871 20030919
INVENTOR(S): Segal, Andrew H.;	US 2004126793 A1 20040701
Young, Elihu	US 2003-666885 20030919
PATENT ASSIGNEE(S): Genitrix, LLC,	US 2004126357 A1 20040701
USA	US 2003-666886 20030919
SOURCE: PCT Int. Appl., 265 pp.	US 2004142889 A1 20040722
CODEN: PIXXD2	US 2003-666898 20030919
DOCUMENT TYPE: Patent	US 2004151728 A1 20040805
LANGUAGE: English	US 2003-666834 20030919
FAMILY ACC. NUM. COUNT: 2	US 2004170960 A1 20040902
PATENT INFORMATION:	US 2003-667193 20030919
	US 2004180389 A1 20040916
PATENT NO. KIND DATE	US 2003-667166 20030919
APPLICATION NO. DATE	US 2004241137 A1 20041202
***************************************	US 2003-666833 20030919
	US 2005064391 A1 20050324
WO 2004018698 A2 20040304	US 2003-668073 20030919
WO 2003-US26072 20030820	PRIORITY APPLN. INFO.: US
W: AE, AG, AL, AM, AT, AU, AZ, BA,	2002-224661 A 20020820
BB, BG, BR, BY, BZ, CA, CH, CN,	US 2002-
CO, CR, CU, CZ, DE, DK, DM, DZ,	404823P P 20020820
EC, EE, ES, FI, GB, GD, GE, GH,	US 2003-
GM, HR, HU, ID, IL, IN, IS, JP, KE,	487407P P 20030715
KG, KP, KR, KZ, LC, LK, LR,	US 2003-645000
LS, LT, LU, LV, MA, MD, MG, MK,	A3 20030820
MN, MW, MX, MZ, NI, NO, NZ, OM,	WO 2003-
PG, PH, PL, PT, RO, RU, SC, SD,	US26072 W 20030820
SE, SG, SK, SL, SY, TJ, TM, TN,	AB The present invention provides a fusion
TR, TT, TZ, UA, UG, US, UZ, VC,	polypeptide which can bind to a
VN, YU, ZA, ZM, ZW	cell surface binding moiety (e.g., a
RW: GH, GM, KE, LS, MW, MZ, SD,	carbohydrate) and server as a ligand
SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY,	for a cell surface polypeptide, as well as
KG, KZ, MD, RU, TJ, TM, AT, BE,	a vector comprising a nucleic
BG, CH, CY, CZ, DE, DK, EE, ES,	acid encoding for such a fusion
FI, FR, GB, GR, HU, IE, IT, LU, MC,	polypeptide, and a host cell comprising
NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN,	such nucleic acid. The lectin is collectin,
GQ, GW, ML, MR, NE, SN, TD, TG	galectin, C-type lectin or
US 2004039156 A1 20040226	glycoprotein; and the cell surface protein is cytokine receptor, CD40,
US 2002-224661 20020820	adhesion mol., defensin receptor, heat
CA 2496384 AA 20040304 CA	shock protein receptor, T cell
2003-2496384 20030820	costimulatory mol., counterreceptor of T
AU 2003265523 A1 20040311	cell costimulatory mol., or
AU 2003-265523 20030820	opsonin receptor. The present invention
US 2004091503 A1 20040513	also provides a compn. comprising
US 2003-645000 20030820	an antigen bearing target and such a
EP 1573047 A2 20050914 EP	fusion polypeptide, as well as a
2003-793170 20030820	compn. comprising a virus or a cell and
R: AT, BE, CH, DE, DK, ES, FR, GB,	such a fusion polypeptide. The
GR, IT, LI, LU, NL, SE, MC, PT,	antigen is tumor antigen, viral antigen,
IE, SI, LT, LV, FI, RO, MK, CY, AL,	bacterial antigen, fungal
TR, BG, CZ, EE, HU, SK	antigen, parasitic antigen, prion antigen,
JP 2006517512 T2 20060727 JP	or autoimmune disease antigen.
2004-531131 20030820	The present invention further relates to a
	method of modulating an immune

response in an animal using such compns. or vaccines.

L1 ANSWER 3 OF 13 CAPLUS COPYRIGHT 2006 ACS on STN

2003:550998 ACCESSION NUMBER:

CAPLUS <<LOGINID::20061019>>

DOCUMENT NUMBER:

139:99846 Vaccination with fusion

TITLE: opsonins targeting

antigen-presenting cells

INVENTOR(S):

Segal, Andrew

PATENT ASSIGNEE(S): USA

U.S. Pat. Appl. Publ.,

SOURCE: 952 pp., Cont.-in-part of U.S.

Ser. No. 789,922.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 5

PATENT INFORMATION:

PATENT NO. APPLICATION NO.

KIND DATE

DATE

A1 20030717 US 2003133942

US 2002-262828

20021002

US 2001031264

A1 20011018

20010221 US 2001-789922

PRIORITY APPLN. INFO.:

US

2001-789922

A2 20010221 US 1996-

11047P P 19960125

US 1998-7711

A2 19980115

AB The author discloses the enhancement of immune responses induced by

in-frame fusion of an antigen with a

binding domain of an opsonin targeting antigen-presenting cells. In one

example, DNA immunization with a chimeric construct of chicken lysozyme

and the .alpha.-chain of

complement C3b increased the IgG1 response over that elicited by a

recombinant lysozyme construct alone.

L1 ANSWER 4 OF 13 CAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2002:716866

CAPLUS <<LOGINID::20061019>>

DOCUMENT NUMBER:

137:231362 SOURCE:

TITLE:

Opsonin fusion proteins

for modulation of the immune

response

INVENTOR(S):

Segal, Andrew

PATENT ASSIGNEE(S): USA

SOURCE:

U.S. Pat. Appl. Publ.,

29 pp., Cont.-in-part of U.S.

6,224,870.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

Enalish

LANGUAGE:

FAMILY ACC. NUM. COUNT: 5 PATENT INFORMATION:

PATENT NO.

KIND DATE

APPLICATION NO. DATE

US 2002131974

A1 20020919

US 2001-790317

20010221

US 6632436

B2 20031014

US 6224870

B1 20010501 US

1998-7711 19980115 PRIORITY APPLN. INFO.:

US

1996-11047P P 19960125

A2 19980115

US 1997-788143

US 1998-7711

B2 19970124

AB The authors discloses the application of

in-frame translation fusion of an

antigen with an APC binding domain of

an opsonin to form a mol., which on

administration, modulates an immune

response to the antigen. In one

example, the IgG1 response was shown

to be enhanced for a construct of

lysozyme and a fragment of the C3b

.alpha.-chain.

L1 ANSWER 5 OF 13 CAPLUS

COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2002:368513

CAPLUS <<LOGINID::20061019>>

DOCUMENT NUMBER:

TITLE:

136:380110 Apolipoprotein A analogs

capable of forming HDL and

with extended serum half-

lives and stronger binding to

cubilin for treatment of

cardiovascular disease

INVENTOR(S):

Graversen, Jonas;

Moestrup, Soren PATENT ASSIGNEE(S):

Aps, Den.

Proteopharma

PCT Int. Appl., 113 pp. CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 1	DK 2001-57
PATENT INFORMATION:	A 20010115
	US 2001-
PATENT NO. KIND DATE	264022P P 20010126
	WO 2001-
APPLICATION NO. DATE	
	DK739 W 20011109
** ***********	US 2001-987107
WO 2002038609 A2 20020516	A3 20011113
WO 2001-DK739 20011109	AB The invention relates to an
WO 2002038609 A3 20020926	apolipoprotein construct, an apolipoprotein
W: AE, AG, AL, AM, AT, AU, AZ, BA,	construct for use as a medicament, a
BB, BG, BR, BY, BZ, CA, CH, CN,	nucleic acid sequence encoding the
CO, CR, CU, CZ, DE, DK, DM, DZ,	apolipoprotein construct, a vector
	comprising the nucleic acid sequence, a
EC, EE, ES, FI, GB, GD, GE, GH,	•
GM, HR, HU, ID, IL, IN, IS, JP, KE,	method for producing the apolipoprotein
KG, KP, KR, KZ, LC, LK, LR,	construct, and use of the
LS, LT, LU, LV, MA, MD, MG, MK,	apolipoprotein construct for the prepn. of
MN, MW, MX, MZ, NO, NZ, OM, PH,	apharmaceutical compn.
PL, PT, RO, RU, SD, SE, SG, SI,	Specifically, analogs and fusion proteins
SK, SL, TJ, TM, TR, TT, TZ, UA,	of apolipoprotein AI are
UG, US, UZ, VN, YU, ZA, ZW	described. The presented data
RW: GH, GM, KE, LS, MW, MZ, SD,	document that the constructs according to
SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,	the invention are capable of binding
DE, DK, ES, FI, FR, GB, GR, IE, IT,	
· · · · · · · · · · · · · · · · · · ·	lipids, are capable of binding
LU, MC, NL, PT, SE, TR, BF,	cubilin, which is a strong Apo Al receptor,
BJ, CF, CG, CI, CM, GA, GN, GQ,	stronger than native Apo A-I
GW, ML, MR, NE, SN, TD, TG	and that the plasma half life of the
CA 2428114 AA 20020516 CA	constructs is at least tripled
2001-2428114 20011109°	compared to native Apo A-I. Together
AU 2002013843 A5 20020521	these data document that the
AU 2002-13843 20011109	constructs according to the invention are
AU 2002213843 A2 20020521	strong candidates for treatment
AU 2002-213843 20011109	of cardiovascular diseases.
BR 2001015257 A 20030812	
BR 2001-15257 20011109	L1 ANSWER 6 OF 13 CAPLUS
EP 1335938 A2 20030820 EP	COPYRIGHT 2006 ACS on STN
2001-982197 20011109	ACCESSION NUMBER: 2002:10696
	CAPLUS < <loginid::20061019>></loginid::20061019>
R: AT, BE, CH, DE, DK, ES, FR, GB,	
GR, IT, LI, LU, NL, SE, MC, PT,	DOCUMENT NUMBER: 136:68702
IE, SI, LT, LV, FI, RO, MK, CY, AL,	TITLE: Analysis of CD154
TR	oligomerization on CD40 signaling
JP 2004522424 T2 20040729 JP	using CD154- ***collectin***
2002-541940 20011109	***fusion***
US 2002156007 A1 20021024	protein
US 2001-987107 20011113	INVENTOR(S): Al-Shamkhani,
US 6897039 B2 20050524	Aymen; Glennie, Martin
NO 2003002101 A 20030708	PATENT ASSIGNEE(S): Cancer
NO 2003-2101 20030509	Research Ventures Limited, UK
ZA 2003004486 A 20040909 ZA	SOURCE: PCT Int. Appl., 63 pp.
2003-4486 20030609	CODEN: PIXXD2
US 2005096277 A1 20050505	DOCUMENT TYPE: Patent
US 2004-17037 20041221	LANGUAGE: English
US 2005142639 A1 20050630	FAMILY ACC. NUM. COUNT: 1
US 2004-17059 20041221	PATENT INFORMATION:
DDIODITY ADDI N. INICO : DV	

DK

PRIORITY APPLN. INFO.:

2000-1682 A 20001110

PATENT NO. KIND DATE more potent than trimeric CD154 in APPLICATION NO. DATE inducing B cell proliferation. Multimeric fusion protein SP-D-CD154 also induced higher levels of WO 2002000893 A1 20020103 expression of ICAM-1 and CD86, WO 2001-GB2810 20010625 compared to those of trimeric CD154. W: AE, AG, AL, AM, AT, AU, AZ, BA, SP-D-CD154 can potentially bind to 12 BB, BG, BR, BY, BZ, CA, CH, CN, CD40 mol., compared to three mols. CO, CR, CU, CZ, DE, DK, DM, DZ, with trimeric CD154, implying that the EC, EE, ES, FI, GB, GD, GE, GH, extent of receptor oligomerization GM, HR, HU, ID, IL, IN, IS, JP, KE, may influence the signals generated by KG, KP, KR, KZ, LC, LK, LR, CD40. REFERENCE COUNT: LS, LT, LU, LV, MA, MD, MG, MK, THERE 4 MN, MW, MX, MZ, NO, NZ, PL, PT, ARE 4 CITED REFERENCES AVAILABLE FOR THIS RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, RECORD. ALL UZ, VN, YU, ZA, ZW, AM, AZ, BY, CITATIONS AVAILABLE IN THE RE KG, KZ, MD, RU, TJ, TM **FORMAT** RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, L1 ANSWER 7 OF 13 CAPLUS DE, DK, ES, FI, FR, GB, GR, IE, IT, COPYRIGHT 2006 ACS on STN LU, MC, NL, PT, SE, TR, BF, ACCESSION NUMBER: 2001:435124 BJ, CF, CG, CI, CM, GA, GN, GW, CAPLUS <<LOGINID::20061019>> 135:45182 ML, MR, NE, SN, TD, TG DOCUMENT NUMBER: CA 2414342 AA 20020103 CA TITLE: Multimeric forms of TNF 2001-2414342 20010625 superfamily ligands EP 1297160 A1 20030402 EP INVENTOR(S): Kornbluth, Richard 2001-945468 20010625 R: AT, BE, CH, DE, DK, ES, FR, GB, PATENT ASSIGNEE(S): USA GR, IT, LI, LU, NL, SE, MC, PT, SOURCE: PCT Int. Appl., 73 pp. IE, SI, LT, LV, FI, RO, MK, CY, AL, CODEN: PIXXD2 TR DOCUMENT TYPE: Patent US 2004047873 A1 20040311 LANGUAGE: English US 2003-312374 20031010 FAMILY ACC. NUM. COUNT: 1 PRIORITY APPLN. INFO.: GB PATENT INFORMATION: 2000-15426 A 20000624 WO 2001-PATENT NO. KIND DATE W 20010625 GB2810 APPLICATION NO. DATE AB The invention provides a protein framework which allows active polypeptides e.g. ligands or antigens to WO 2001042298 A1 20010614 be displayed at increased concn. WO 2000-US7380 20000320 The inventors show that the lectin binding W: AU, CA, JP domains of collectins can be RW: AT, BE, CH, CY, DE, DK, ES, FI, replaced by a polypeptide of interest and FR, GB, GR, IE, IT, LU, MC, NL, that polypeptide can be PT, SE AA 20010614 CA multimerised by the framework of the CA 2393659 2000-2393659 collectin and as a result displayed 20000320 in greater no. on a single structure. The EP 1235853 A1 20020904 inventors show that the 2000-919485 20000320 R: AT, BE, CH, DE, DK, ES, FR, GB, activity of polypeptides such as those of the TNF superfamily are GR, IT, LI, LU, NL, SE, MC, PT, significantly enhanced when displayed in IE, FI, CY this way. The invention US 2005158831 A1 20050721 demonstrated that multimeric fusion US 2005-87348 20050322 protein SP-D-CD154 was about 8 fold

PRIORITY APPLN. INFO.: US 1999-454223 A 19991209 US 1998-P 19981209 111471P WO 2000-US7380 W 20000320 AB A method for constructing stable bioactive fusion proteins of the difficult to express tumor necrosis factor superfamily (TNFSF), and particularly members CD40L (CD154) and RANKL/TRANCE, with collectins, particularly pulmonary surfactant protein D (SPD) is described. Single trimers of these proteins lack the full stimulatory efficacy of the natural membrane forms of these proteins in many cases. The multimeric nature of these sol. fusion proteins enables them to engage multiple receptors on the responding cells, thereby, mimicking the effects of the membrane forms of these ligands. For CD40L-SPD, the resulting protein stimulates B cells, macrophages, and dendritic cells, indicating its potential usefulness as a vaccine adjuvant. The large size of these fusion proteins makes them less likely to diffuse into the circulation, thereby limiting their potential systemic toxicity. This property may be esp. useful when these proteins are injected locally as a vaccine adjuvant or tumor immunotherapy agent to prevent them from diffusing away. In addn., these and other TNFSF-***collecting*** ***fusion*** proteins present new possibilities for the expression of highly active, multimeric, sol. TNFSF members. REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE **FORMAT** L1 ANSWER 8 OF 13 CAPLUS COPYRIGHT 2006 ACS on STN 2001:310484 ACCESSION NUMBER:

CAPLUS <<LOGINID::20061019>>

comprising antigens fused with

134:325200

Vaccine compns.

DOCUMENT NUMBER:

TITLE:

antigen-presenting cellbinding domains of opsonins INVENTOR(S): Segal, Andrew H. PATENT ASSIGNEE(S): Genitrix, Ltd., USA SOURCE: · U.S., 21 pp., Cont.-inpart of U.S. Ser. No. 788,143, abandoned. CODEN: USXXAM DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: 5 PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE US 6224870 B1 20010501 US 19980115 1998-7711 WO 9936507 A1 19990722 19990115 WO 1999-US894 W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG AU 9922301 A1 19990802 AU 1999-22301 19990115 US 2001031264 A1 20011018 US 2001-789922 20010221 US 2002131974 A1 20020919 US 2001-790317 20010221 B2 20031014 US 6632436 PRIORITY APPLN. INFO.: US 1997-788143 B2 19970124 US 1996-11047P P 19960125 US 1998-7711 A 19980115 WO 1999-**US894** W 19990115

AB The invention provides compns. and methods for modulating immune responses

in subjects. The invention is based, at RECORD. ALL least in part, on the discovery CITATIONS AVAILABLE IN THE RE that an in-frame translation fusion of an **FORMAT** antigen with an APC binding domain of an opsonin forms a mol., i.e., a L1 ANSWER 10 OF 13 CAPLUS fusion polypeptide, which when COPYRIGHT 2006 ACS on STN administered to a subject modulates an ACCESSION NUMBER: 1991:459229 CAPLUS <<LOGINID::20061019>> immune response to the antigen. REFERENCE COUNT: DOCUMENT NUMBER: 115:59229 41 THERE Methods and systems for ARE 41 CITED REFERENCES AVAILABLE TITLE: FOR THIS generating and RECORD. ALL ***collecting*** CITATIONS AVAILABLE IN THE RE ***fusion*** fuel material **FORMAT** INVENTOR(S): Lautzenhiser, Theodore V.; Eisner, Melvin L1 ANSWER 9 OF 13 CAPLUS PATENT ASSIGNEE(S): Amoco Corp., COPYRIGHT 2006 ACS on STN USA ACCESSION NUMBER: 2000:753729 SOURCE: Can. Pat. Appl., 17 pp. CAPLUS <<LOGINID::20061019>> CODEN: CPXXEB DOCUMENT NUMBER: 134:351860 DOCUMENT TYPE: Patent TITLE: Development of chimeric LANGUAGE: English collectins with enhanced FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION: activity against influenza A virus KIND DATE AUTHOR(S): Hartshorn, Kevan L.; PATENT NO. APPLICATION NO. White, Mitchell R.; Alan, R.; DATE Ezekowitz, B.; Sastry, Kedarnath; Crouch, Erika CA 2005641 CORPORATE SOURCE: Boston AA 19901212 CA 1989-2005641 University School of Medicine, Boston, MA, 19891215 USA PRIORITY APPLN. INFO.: US SOURCE: Advances in 1989-364936 A 19890612 Experimental Medicine and Biology (2000), AB Methods and systems are described for 479(Biology and Pathology the generation and collection of T. of Innate Immunity In particular, T is generated at a reducing electrode of a Galvanic cell Mechanisms), 49-59 and thereafter biased so as to migrate to CODEN: AEMBAP; ISSN: 0065-2598 a selected surface of the PUBLISHER: reducing electrode. T which is migrated Kluwer Academic/Plenum Publishers to the selected surface and DOCUMENT TYPE: Journal; General coalesced thereon is then collected. Review LANGUAGE: L1 ANSWER 11 OF 13 USPATFULL on English AB A review with 30 refs. Topics STN discussed include the functional ACCESSION NUMBER: 2005:183444 USPATFULL <<LOGINID::20061019>> significance of variations in carbohydrate binding specificity and TITLE: Multimeric fusion proteins quaternary structure of collectins with of TNF superfamily ligands respect to influenza A virus INVENTOR(S): Kornbluth, Richard infection; and construction of collectin S., La Jolla, CA, UNITED STATES chimeras to det. the contribution of specific domains to antiviral and NUMBER KIND opsonic activities. DATE REFERENCE COUNT: 30 THERE ARE 30 CITED REFERENCES AVAILABLE PATENT INFORMATION: US

A1 20050721

2005158831

FOR THIS

APPLICATION INFO.: US 2005-87348 A1 20050322 (11) RELATED APPLN. INFO.: Continuation of Ser. No. US 1999-454223, filed on 9 Dec 1999, PENDING

NUMBER . DATE

PRIORITY INFORMATION: US 1998-111471P 19981209 (60) DOCUMENT TYPE: Utility FILE SEGMENT: APPLICATION LEGAL REPRESENTATIVE: Lisa A. Haile. J.D., Ph.D., DLA PIPER RUDNICK GRAY CARY

US LLP, Suite 1100, 4365

Executive Drive, San Diego,

CA, 92121-2133, US

NUMBER OF CLAIMS: **EXEMPLARY CLAIM:**

24 1-15

NUMBER OF DRAWINGS: 7 Drawing

Page(s)

LINE COUNT:

1657

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

AB A method for constructing stable bioactive fusion proteins of the difficult to express turn or necrosis factor superfamily (TNFSF), and particularly members CD40L (CD 154) and RANKL/TRANCE, with collectins, particularly pulmonary surfactant protein D (SPD) is described. Single

trimers of these proteins lack the full stimulatory efficacy of the

natural membrane forms of these proteins in many cases. The multimeric nature of these soluble fusion proteins enables them to engage multiple

receptors on the responding cells, thereby, mimicking the effects of the membrane forms of these ligands. For CD40L-SPD, the resulting protein

stimulates B cells, macrophages, and dendritic cells, indicating its

potential usefulness as a vaccine adjuvant. The large size of these

fusion proteins makes them less likely to diffuse into the circulation,

thereby limiting their potential systemic toxicity. This property may be

especially useful when these proteins are injected locally as a vaccine

adjuvant or tumor immunotherapy agent to prevent them from diffusing

away. In addition, these and other TNFSF- ***collectin***

fusion proteins present new possibilities for the expression of

highly active, multimeric, soluble TNFSF members.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 12 OF 13 USPATFULL on STN

ACCESSION NUMBER: 2004:292713 USPATFULL <<LOGINID::20061019>>

TITLE: Methods and

compositions for treating ocular disease INVENTOR(S): Fleiszig, Suzanne

M.J., Oakland, CA, UNITED STATES Evans, David J., Oakland,

CA, UNITED STATES

Sack, Robert A., Brookhaven, NY, UNITED STATES

> NUMBER KIND

DATE

PATENT INFORMATION: US

A1 20041118 2004229802 APPLICATION INFO.: US 2004-823819

A1 20040414 (10)

NUMBER DATE

PRIORITY INFORMATION: US 2003-

462913P 20030415 (60)

DOCUMENT TYPE:

Utility

FILE SEGMENT: APPLICATION LEGAL REPRESENTATIVE: WOODCOCK WASHBURN LLP, ONE LIBERTY PLACE, 46TH FLOOR,

1650 MARKET STREET,

PHILADELPHIA, PA, 19103

NUMBER OF CLAIMS: 22

EXEMPLARY CLAIM:

NUMBER OF DRAWINGS: 7 Drawing

Page(s)

LINE COUNT:

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

1624

The use of collectins and/or surfactant proteins for the treatment and prevention of ocular disease.

CAS INDEXING IS AVAILABLE FOR THIS PATENT.

L1 ANSWER 13 OF 13 USPATFULL on STN

ACCESSION NUMBER: 92:82537 USPATFULL <<LOGINID::20061019>>

TITLE:

Storage ring fusion energy

generator

INVENTOR(S): Russell, Joseph A., 600 Star Rte., Lompoc, CA, United

States 93436

NUMBER KIND

DATE

PATENT INFORMATION: US 5152955

19921006

APPLICATION INFO.: US 1990-566054

19900809 (7)

DOCUMENT TYPE: Utility FILE SEGMENT: Granted

PRIMARY EXAMINER: Wasil, Daniel D. LEGAL REPRESENTATIVE: Wedemeyer,

Lowell R.

NUMBER OF CLAIMS: 8

EXEMPLARY CLAIM: 6

NUMBER OF DRAWINGS: 9 Drawing

Figure(s); 3 Drawing Page(s) LINE COUNT: 967

AB This invention relates to adaptation of

intersecting storage rings, of

the same type used in high energy nuclear physics research, for power

generation. The device is optimized for

lower-energy beam paricles and

higher beam current, adapted with a

reaction chamber at the intersection of the rings to collect released fusion

energy for conversion to

electricity, and equipped with means to recapture scattered accelerated

particles and reintegrate them into the

focused beams for recirculation

through the reaction chamber. The preferred beam particles, deuterium

and tritium, are accelerated and injected into and focused by the

storage rings, to collide nearly head on in the reaction chamber.

Non-colliding, accelerated beam particles are conserved by recovery, correction and recirculation, requiring relatively small amounts of input energy to maintain acceleration

and focus of the beams, and thus remain energized for another collision attempt. Grid devices intercept

scattered particles and recapture some of them for recirculation. Only those beam particles which scatter so widely as to evade recapture and those which actually react to produce thermonuclear fusion must be replaced and accelerated up to the energy sufficient to cause fusion.